

Landscape-Genetic Analysis of Population Structure in the Texas Gray Fox Oral Rabies Vaccination Zone

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ABSTRACT In west-central Texas, USA, abatement efforts for the gray fox (*Urocyon cinereoargenteus*) rabies epizootic illustrate the difficulties inherent in large-scale management of wildlife disease. The rabies epizootic has been managed through a cooperative oral rabies vaccination program (ORV) since 1996. Millions of edible baits containing a rabies vaccine have been distributed annually in a 16-km to 24-km zone around the perimeter of the epizootic, which encompasses a geographic area $>4 \times 10^5$ km². The ORV program successfully halted expansion of the epizootic into metropolitan areas but has not achieved the ultimate goal of eradication. Rabies activity in gray fox continues to occur periodically outside the ORV zone, preventing ORV zone contraction and dissipation of the epizootic. We employed a landscape-genetic approach to assess gray fox population structure and dispersal in the affected area, with the aim of assisting rabies management efforts. No unique genetic clusters or population boundaries were detected. Instead, foxes were weakly structured over the entire region in an isolation by distance pattern. Local subpopulations appeared to be genetically non-independent over distances >30 km, implying that long-distance movements or dispersal may have been common in the region. We concluded that gray foxes in west-central Texas have a high potential for long-distance rabies virus trafficking. Thus, a 16-km to 24-km ORV zone may be too narrow to contain the fox rabies epizootic. Continued expansion of the ORV zone, although costly, may be critical to the long-term goal of eliminating the Texas fox rabies virus variant from the United States. (JOURNAL OF WILDLIFE MANAGEMENT 73(8):1292–1299; 2009)

DOI: 10.2193/2008-336

KEY WORDS gray fox, landscape genetics, oral rabies vaccination, population structure, spatial autocorrelation, *Urocyon cinereoargenteus*.

The effective control and abatement of animal disease outbreaks is an emerging wildlife management problem that is expected to increase in coming decades with the rise in globalization, urban encroachment, and spread of invasive species (DeYoung 2007). The highly publicized recent occurrences of foot-and-mouth disease in the United Kingdom and chronic wasting disease and bovine tuberculosis in the United States underscore the difficulty of spatially extensive management. Although rabies receives less media attention, the occurrence and management of animal rabies poses a significant problem. Rabies is a worldwide public health threat, causing an estimated 30,000–50,000 human deaths (primarily in third world countries) each year, with many millions of dollars spent on treatment and prevention (Real et al. 2005). In the United States, animal rabies generally occurs in free-ranging species of mammals, often small carnivores and bats, where genetically distinct rabies strains are present in discrete geographical areas. For instance, approximately 92% of rabies cases in the United States during 2004 were in wild animals (Krebs et al. 2005). Rabies transmission in wild populations occurs primarily among conspecifics and within defined geographic regions; the rate of interspecific

infection is generally low. Once the virus becomes established, rabies outbreaks may achieve long-term persistence (e.g., several decades or more) within geographic regions (Real et al. 2005).

A case cluster of rabies in gray fox (*Urocyon cinereoargenteus*) in eastern Texas, USA, in 1946 signaled the beginning of a gray fox rabies epizootic (Texas fox rabies virus variant [TF]) in the state (Texas Department of Health 2003), and in 1988 a major epizootic was noted (Sidwa et al. 2005). By 1994, this epizootic, and a simultaneous epizootic in coyotes (*Canis latrans*) and domestic dogs in south Texas (domestic dog-coyote rabies virus variant [DDC]), were declared as state health emergencies (United States Department of Agriculture [USDA] 2002). To combat the potential public health risks, oral rabies vaccination (ORV) programs were initiated for wildlife in southern Texas for the DDC variant in 1995 and in west-central Texas for the TF variant in 1996 (Sidwa et al. 2005), with the ultimate goal of eliminating these strains of rabies virus from the United States (USDA 2001). The ORV programs used a recombinant vaccinia-rabies vaccine (Raboral V-RG®; Rhone Merieux, Inc., Athens, GA) developed for the oral vaccination of select wildlife species against rabies virus. The general vaccination strategy was to distribute edible baits containing the rabies vaccine, creating a geographic zone of vaccinated individuals

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at the perimeter of the epizootic (Sidwa et al. 2005). Animals consuming the baits become immunized and the enzootic is disrupted when a sufficient portion of the population is vaccinated (i.e., the herd immunity concept; Fox et al. 1995).

Overall, ORV programs have yielded exceptional success with the use of Raboral V-RG (Sidwa et al. 2005). For instance, the DDC variant was effectively eliminated from south Texas by 2003 (Sidwa et al. 2005). The gray fox ORV program was successful in halting expansion of the TF variant, but rabies activity in Texas gray foxes continued to occur over a large geographic expanse of west-central Texas (Texas Department of Health 2003, Sidwa et al. 2005). The gray fox ORV program attempted to halt the eastern and northward expansion of the virus toward metropolitan areas of central Texas (including San Antonio) and the large red fox (*Vulpes vulpes*) populations in eastern Texas by creating a 16-km to 24-km wide vaccination zone, which was adjusted annually depending on resource availability and distribution of positive individuals (Sidwa et al. 2005). It was hypothesized that this strategic ORV zone would eventually reduce and encircle the affected area, with the ultimate goal of eliminating the TF variant from the United States.

Each year since the inception of the program, >1 million ORV baits have been distributed over thousands of square kilometers in west-central Texas. In recent years, the gray fox ORV zone in Texas has been large, extending from the border of Mexico to west-central Texas (Fig. 1). To date, rabies abatement of the affected area has not been achieved, as demonstrated by the periodic detection of rabid animals outside, but near the perimeter of, the gray fox ORV zone during evaluations following several annual bait dispersal campaigns (e.g., 1999, 2002, 2004, 2007; G. Moore, Texas Department of State Health Services, unpublished). The disparate success of the coyote versus the fox ORV may be partly due to the extensive geographic area of the fox ORV and the difficulty in allocating sufficient resources to blanket the entire affected area. However, the geographic distribution of rabies-infected mammals in other regions of the United States appears to be influenced in part by terrain features that affect animal movements (Childs et al. 2000, 2001; Smith et al. 2002). Thus, ecological information on gray fox movements and population structure in west-central Texas could improve the success of the gray fox ORV program. Unfortunately, much information pertaining to general ecology of gray fox in west-central Texas remains undetermined.

Gray foxes are distributed statewide in Texas (Schmidly 1994), though some have suggested that gray foxes may be restricted to habitats where they are not excluded through behavioral interactions with sympatric coyotes and bobcats (*Lynx rufus*; MacDonald and Sillero-Zubiri 2004). Gray foxes exist in family units composed of an adult pair and their juvenile offspring (Schmidly 1994); juveniles may remain in their natal range until January–February the following year (Trapp and Hallberg 1975). Gray foxes are social and the (presumably monogamous) adult pairs have been observed to travel together within their home ranges (Chamberlain and Leopold 2000). Tagging and telemetry

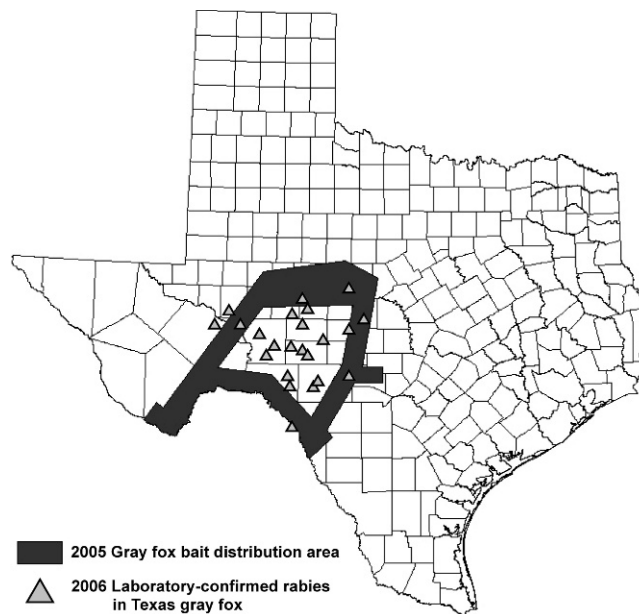


Figure 1. Oral rabies vaccination zone for gray fox in west-central Texas, USA, during 2005 and location of laboratory-confirmed cases of gray fox rabies in 2006. The zone and sample collection area encompassed all or parts of 33 counties.

studies have reported occasional long-distance movements (20–80 km; Sheldon 1953, B. Mesenbrink, USDA, Wildlife Services, National Wildlife Research Center, unpublished data), but the frequency, average distance, and the degree to which long-distance dispersers integrate into the population are unknown.

Information on the population structure and movement patterns of gray fox would allow for better use of the limited resources available for managing animal rabies in this geographically expansive area. Traditional wildlife studies, employing tagging and radiotelemetry, are informative but limited by expense and constraints on sample size. Recent advances in molecular techniques have made large-scale genetic analyses of wildlife populations feasible (DeYoung and Honeycutt 2005), offering powerful new tools for achieving insights into animal behavior and movements (DeYoung 2007). An increasing number of studies have begun to use genetic information to assist large-scale applied management (e.g., Root et al. 2003, Hampton et al. 2004), an analytical approach that is now termed *landscape genetics* (Manel et al. 2003).

We conducted a series of population genetic analyses aimed at assisting gray fox ORV strategies in western Texas. The specific objective of our study was to estimate the degree and spatial extent of population structuring in the ORV region. If structuring was detected, we would attempt to identify landscape features influencing population structure.

STUDY AREA

Our study focused on the area inside and around the perimeter of the Texas gray fox ORV zone in west-central Texas. The geographic extent of the ORV zone has varied in geographic extent since its inception in 1996. During study

collection, the specific area encompassed 33 counties, a geographic area of about $4 \times 10^5 \text{ km}^2$ (Fig. 1). The ORV zone included much of the Edwards Plateau, a semiarid region comprised mostly of rangelands characterized by rocky plains interspersed with hills and steep drainages (Hatch et al. 1990).

METHODS

DNA Extraction and Amplification

We obtained tissue biopsies (ear punch or muscle) from gray fox during 2005–2006 (Fig. 1). We trapped or collected gray foxes during evaluations of annual ORV bait dispersion (Sidwa et al. 2005), ongoing research projects, road kills, and wildlife damage management activities. For each tissue sample collected, we recorded corresponding Global Positioning System coordinates of the collection site along with the sex and approximate age (juv or ad). We placed tissues in cryovials containing 1 mL of 70% ethanol and maintained them at -20°C until DNA extractions.

We extracted DNA using a commercial kit (DNeasy tissue kit; Qiagen Genomics, Valencia, CA) and used the polymerase chain reaction (PCR) to amplify 5 microsatellite loci (GF-02, GF-07, GF-09, GF-12, and GF-14) from a panel designed for gray fox (Weston et al. 2004). We amplified microsatellite loci in 25- μL reaction volumes containing 12.5 μL Amplitaq Gold PCR Master Mix (a premixed solution of thermal-stable DNA polymerase, dNTP, Mg^{++} , and buffer; Applied Biosystems, Foster City, CA), 10 picomoles (pmol) each primer, and 10–50 ng DNA. Thermal cycling conditions are described in Weston et al. (2004). We combined the PCR products (3 μL each reaction) for each individual and applied 1 μL of the mixture to a denaturing formamide and size standard mix (Hi-Di Formamide, ROX 500, Applied Biosystems). We then loaded the PCR product mixtures onto an ABI 3130 \times DNA sequencer (Applied Biosystems) for separation and detection. We quantified fragment sizes and resolved into multilocus genotypes using *GENEMAPPER* 4.0 software (Applied Biosystems), followed by visual inspection and verification.

We sequenced a portion of the hypervariable mtDNA control region (Domain I) using primers LGL 283F, 5'-TACACTGGTCTTGTAAC-3', and LGL 1115R, 5'-ATGACCCTGAAGAARGAACCAG-3' (Harlin-Cognato et al. 2006). We amplified the control region sequences in 25- μL reaction volumes containing 12.5 μL Amplitaq Gold PCR Master Mix (Applied Biosystems), 10 pmol of each primer, and 10–50 ng DNA. Reaction conditions consisted of an initial denaturation at 94°C for 12 minutes followed by 32 cycles of 94°C for 50 seconds, 61°C for 60 seconds, 72°C for 2 minutes, with a final extension at 72°C for 30 minutes. We electrophoresed PCR products on 1% agarose gels containing ethidium bromide and viewed under ultraviolet light to verify successful amplification. We purified products from successful reactions using an enzymatic method (ExoSAP-IT; USB Corporation, Wilmington, MD), then used the purified products as templates for sequencing reactions using the BigDye Terminator Cycle Sequencing kit v1.1 (Applied Biosystems). We removed unincorporated dye terminators

(DyeEx 2.0 spin kit, Qiagen) and sequenced each sample in both directions on an ABI 3130 \times automated DNA sequencer (Applied Biosystems).

Data Analysis

We performed 3 analyses aimed at quantifying the number of discrete groups and the spatial extent of population structure in the gray fox ORV zone. First, we used the computer program *FSTAT* 2.9.3 (Goudet 2001) to test for both Hardy–Weinberg equilibrium and for genetic structure over the entire data set based on F_{ST} (Weir and Cockerham 1984). We tested for significant departure of global F_{ST} from zero by jackknifing over loci, while significance of departure from Hardy–Weinberg expectations was assessed by 1,000 randomizations of alleles among individuals. We performed a locus-by-locus analysis of molecular variance (AMOVA; Weir and Cockerham 1984, Excoffier et al. 1992, Weir 1996) and exact tests of population differentiation using the computer program *ARLEQUIN* 3.1 (Excoffier et al. 2005), where gray fox samples collected within counties were pooled and each county was treated as a discrete population. We only conducted AMOVA and exact tests among counties with an arbitrary minimum of 13 gray fox samples (aimed at gaining reasonable estimates of allele frequencies). The AMOVA analysis partitions genetic variation among populations (counties) and individuals, while the exact tests of population differentiation test the hypothesis of a random distribution of genotypes among populations. Statistical significance of the AMOVA was assessed by 1,023 permutations of genotypes among counties; statistical significance of the exact tests was assessed with the Markov chain procedure of Guo and Thompson (1992).

Next, we used a Bayesian clustering algorithm (Pritchard et al. 2000) implemented in the computer program *STRUCTURE* 2.1 (<http://pritch.bsd.uchicago.edu/software.html>, accessed 31 Aug 2009) to assess the number of discrete genetic clusters (K) present in the ORV zone. The algorithm groups individuals into clusters that minimize Hardy–Weinberg and gametic disequilibrium without regard to the spatial location of the sample, requiring no a priori assumptions about population (cluster) boundaries. We used the admixture model and assumed allele frequencies to be correlated. We performed 100,000 burn-in repetitions (reps) to minimize the effect of the starting configuration, followed by 250,000 reps of data collection. We performed 10 independent runs for each K (assumed no. of genetic clusters) to evaluate consistency of the results. We performed 2–3 additional runs of varying lengths for each K to further evaluate consistency of the results. We used the mean and standard deviation of the estimated log probability of data, $L(K)$, among runs for the same K to assess the most likely number of unique clusters supported by the data.

Finally, we performed an analysis of spatial autocorrelation to investigate the spatial extent of genetic structure in the ORV zone. Spatial autocorrelation quantifies the degree to which individual genotype frequencies are correlated as a function of the Euclidian geographic distance between pairs of individuals, useful for defining the spatial extent of

Table 1. Observed (H_O) and expected heterozygosity (H_E), number of alleles (k), and fixation indices (F_{IS} and F_{ST}) for 5 microsatellite DNA loci amplified in a gray fox population sampled in west-central Texas, USA, during 2004–2005.

Locus	n	H_O	H_E	k	F_{IS}	SE	F_{ST}	SE
GF-02	454	0.75	0.89	23	0.177	0.020*	0.007	0.004
GF-07	461	0.75	0.80	17	0.053	0.035	−0.002	0.004
GF-09	457	0.87	0.85	9	−0.034	0.021	0.007	0.003*
GF-12	369	0.76	0.76	16	0.023	0.029	0.012	0.010
GF-14	467	0.70	0.72	9	0.036	0.036	0.013	0.008
Mean or total		0.75		74	0.051	0.039	0.007	0.003*

* 95% CI does not include zero.

population structuring (Manel et al. 2003). We used Moran's I (Moran 1950, Sokal and Oden 1978) as a measure of autocorrelation because the performance of, and theoretical basis for, Moran's I has been extensively investigated in simulation and empirical studies (Hardy and Vekemans 1999, Epperson 2004). We estimated Moran's I (averaged over loci) for all pairs of individuals separated by distance intervals of 10 km, out to 300 km. We used 10-km intervals to assess autocorrelation on a scale relevant to the width of the annual bait drops in the ORV zone (approx. 16–24 km), the area contained by the ORV zone (approx. 300 km in diam), and the occasional long-distance movements observed in previous studies (≤ 80 km). We tested the statistical significance (2-sided) of Moran's I for each distance class by comparing the observed value versus a null value derived from 1,000 permutations of individual locations. We estimated standard errors of I -values by jackknifing over loci. We performed analyses using the computer program *SPAGEDi* 1.2 (Hardy and Vekemans 2002).

We assembled, aligned, and edited the mtDNA sequences using the computer program *SEQUENCHER* 4.5 (Gene Codes, Ann Arbor, MI). We indexed genetic diversity at the haplotype (H) and nucleotide (π) level using the computer program *DNASP* (Rozas et al. 2003). Haplotype diversity is the probability that any 2 randomly sampled haplotypes are different, while π is average number of nucleotide differences per site (Nei 1987). We estimated F_{ST} among counties (0; Weir and Cockerham 1984, Excoffier et al. 1992) as a measure of population structure based on mtDNA haplotypes using the computer program *ARLEQUIN* 3.1.

RESULTS

We genotyped 469 adult foxes from the 33 counties at 5 microsatellite loci. Sampled foxes had high levels of genetic diversity, in terms of both observed heterozygosity and number of alleles (Table 1). Locus GF-02 had a slight excess of homozygotes, possibly due to the presence of null or non-amplifying alleles at low frequency. Locus GF-12 was problematic to score at times due to unusual patterns of stutter. Altering PCR conditions improved this somewhat, but there remained cases where it was difficult to reliably make allele size-calls for a second allele. In the cases where a second allele appeared to be present but we could not confidently establish a size-call, we followed a conservative strategy of coding the second allele as missing to distinguish this case from a true homozygote. As a result, fewer complete genotypes were obtained for this locus than the other four (Table 1). We

observed slight but statistically significant genetic structure ($F_{ST} = 0.007$) over the entire data set; F_{IS} was positive but was not statistically different from zero (Table 1).

We considered 14 counties for AMOVA and exact tests of differentiation ($n = 359$, range = 13–57, median/county = 20.5). The AMOVA revealed that 94.3% of genetic variation was contained within individuals, 5.0% among individuals within counties, and only 0.7% among counties. The exact tests of differentiation revealed no statistically significant differences among all counties (global P -value > 0.79), or for pairwise tests between counties ($P > 0.10$). The *STRUCTURE* results revealed that the maximal value of $L(K)$ was attained at $K = 1$ (Fig. 2). The $L(K)$ values decreased for each $K > 1$ and became more variable among runs. The proportion of individuals in each inferred cluster for $K > 1$ were evenly distributed (e.g., for $K = 2$, each cluster contained 50% of individuals, for $K = 3$, each cluster contained 33% of individuals, etc.). The maximum values of $L(K)$ never reached a plateau, so there was no need to employ the rate of change metrics recommended by Evanno et al. (2005) for identification of genetic clusters. We concluded that the AMOVA, exact test, and *STRUCTURE* results supported only a single genetic cluster in the ORV zone.

Statistical power for spatial autocorrelation analyses can be indexed by the total number of alleles (k) multiplied by the number of sampled individuals (N); if the product ($k \times N$) is at least several thousand, tests for I have sufficient power (Epperson 2005). Thus, our sample of 469 foxes and 74 alleles ($k \times N = 34,706$) provided more than adequate statistical power to detect departures from equilibrium. Indeed, the autocorrelation coefficients revealed weak but statistically

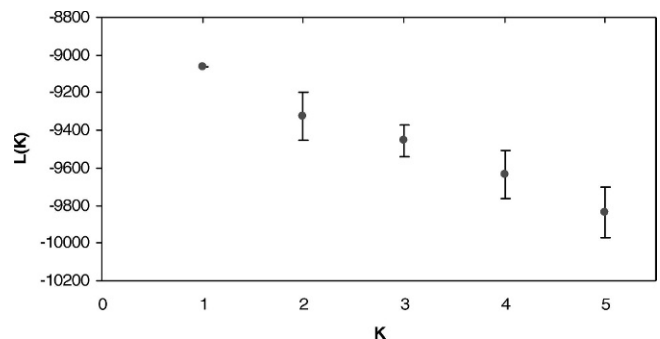


Figure 2. Log probability of data [$L(K)$] as a function of K averaged over 10 independent runs for gray fox in west-central Texas, USA, derived using a Bayesian clustering algorithm implemented in the computer program *STRUCTURE*. The Y-error bars are ± 1 standard deviation and K is the assumed number of genetic clusters.

Table 2. Observed and permuted values of spatial autocorrelation coefficients (Moran's *I*) averaged over 5 microsatellite loci for 469 gray foxes collected during 2004–2005 in the Texas, USA, oral rabies vaccination zone over 30 10-km distance classes.

Class ^a	No. pairs ^b	%	CV ^c	Observed	SE	Permuted	P-value ^d
10	2,155	90.4	1.12	0.021	0.010	2.2×10^{-6}	<0.001***
20	1,443	88.3	0.87	0.013	0.004	3.0×10^{-3}	0.035**
30	2,090	94.7	0.88	0.014	0.003	-9.3×10^{-5}	0.003***
40	2,335	97.9	0.71	-0.002	0.006	-1.4×10^{-4}	0.628
50	2,559	97.0	0.67	0.002	0.003	-5.6×10^{-5}	0.588
60	2,089	94.9	0.80	0.010	0.005	-2.6×10^{-3}	0.055*
70	1,870	97.0	0.93	0.009	0.006	-2.0×10^{-4}	0.075*
80	2,231	94.9	0.83	-3.6×10^{-4}	0.006	-1.5×10^{-4}	0.998
90	2,588	90.8	0.91	-0.002	0.003	-5.8×10^{-5}	0.718
100	2,822	97.4	0.86	0.017	0.001	-1.1×10^{-4}	<0.001***
110	3,468	97.4	0.79	2.4×10^{-4}	0.005	-1.7×10^{-4}	0.910
120	4,078	96.6	0.75	0.007	0.004	-9.6×10^{-5}	0.019**
130	4,030	99.6	0.63	0.005	0.002	-7.2×10^{-5}	0.083*
140	4,479	98.9	0.67	-0.005	0.005	-2.7×10^{-5}	0.147
150	4,632	99.1	0.72	-0.001	0.006	-5.3×10^{-5}	0.654
160	3,975	98.5	0.70	-0.001	0.004	-1.9×10^{-4}	0.826
170	4,082	99.6	0.65	0.003	0.002	-2.7×10^{-5}	0.363
180	3,827	100	0.62	0.003	0.004	-9.2×10^{-5}	0.319
190	4,727	100	0.68	-0.002	0.002	-5.6×10^{-6}	0.562
200	3,844	99.8	0.64	-0.004	0.004	3.6×10^{-5}	0.307
210	4,038	100	0.55	0.001	0.004	1.8×10^{-4}	0.786
220	4,049	100	0.63	-0.005	0.003	3.3×10^{-5}	0.147
230	4,774	100	0.58	-0.006	0.003	-2.6×10^{-4}	0.043**
240	4,466	99.6	0.61	-0.006	0.005	-1.5×10^{-5}	0.043**
250	4,362	99.6	0.63	-0.002	0.005	-2.2×10^{-5}	0.588
260	3,683	94.9	0.67	-0.003	0.002	-1.4×10^{-4}	0.425
270	3,480	94.5	0.65	-0.002	0.003	1.1×10^{-4}	0.614
280	2,987	92.1	0.73	-0.008	0.004	3.4×10^{-5}	0.033**
290	2,594	91.3	0.80	0.004	0.010	-2.6×10^{-4}	0.353
300	2,100	86.6	0.93	0.001	0.005	-2.8×10^{-4}	0.696

^a Upper distance bound (km).

^b No. of pairwise comparisons within each interval.

^c Percentage of sampled individuals participating at least once in the interval and the coefficient of variation of the no. of times each individual is represented.

^d Two-sided P-values: * $P < 0.1$; ** $P < 0.05$; *** $P < 0.01$.

significant positive autocorrelation over the first 3 10-km distance classes (Table 2). Five additional statistically significant, positive *I*-values spanned distance classes 60–130 km, with negative correlations observed at greater spatial distance classes (Table 2), suggesting an isolation by distance pattern.

We sequenced 401 base pairs (397 base pairs excluding alignment gaps) for 25 adult female gray foxes. We resolved the sequences into 7 haplotypes; *H* and π were 0.67 (SD = 0.10) and 0.007 (SD = 0.003), respectively. Tests of genetic structure among haplotypes revealed no statistically significant structure ($F_{ST} = -0.13$, $P = 0.62$) among counties in the ORV zone. No geographic pattern to the distribution of mtDNA haplotypes was apparent upon visual inspection.

DISCUSSION

We detected only weak genetic structure over the spatially extensive gray fox ORV zone in Texas, a conclusion substantiated by several different analytical approaches using both microsatellite and mtDNA markers. There was no evidence for unique genetic clusters, which implies that landscape features have a minimal effect on gray fox population structure in the region. At the very least, no habitat features affected population structuring to the extent that they could be useful to the ORV management. Thus, we were unable to satisfy one of the main goals of the study, the identification of landscape features affecting gray fox

population structure that could be used in management. Instead, gray foxes appeared to be weakly structured as a function of geographic distance between individuals. The weak structuring followed an isolation by distance pattern, where greater than expected autocorrelation was observed at proximate distance classes, and lower than expected observed at extreme distance classes. Our autocorrelation values were comparable to those observed in fine-scale studies of dispersal in plants and small mammals, where distance intervals were measured in tens of meters (e.g., Marquardt and Epperson 2004, Walter and Epperson 2004, Peakall et al. 2005). Although the autocorrelation coefficients appeared low, they were actually quite surprising for distance intervals measured in tens of kilometers and corresponded to theoretical expectations for species with high dispersal; simulation studies indicate that Moran's *I*-values in the range of 0.03–0.06 and 0.0–0.02 for the first distance category in a correlogram reflected Wright's (1946) neighborhood sizes of 100–250 and >250 individuals, respectively (Epperson 2003). Thus, the autocorrelation analyses strongly suggested that population structure in the gray fox ORV zone extended to ≤ 30 km, and possibly farther. A simple and conservative interpretation of the results would be that gray fox populations within 30 km are genetically non-independent (Diniz-Filho and Telles 2002).

Comparable studies of gray fox population structure have not been reported from other parts of the species' range, with the exception of congeneric island fox (*U. littoralis*) in the Channel Islands of California, USA, where foxes display dramatic population genetic structuring among islands due to restricted over-water dispersal (Goldstein et al. 1999). Studies of widely distributed species of carnivores have revealed that long-distance dispersal can result in weak genetic structuring at spatial scales many times larger than the gray fox ORV zone (Schwartz et al. 2002). In many canid and felid species, structuring at such large spatial scales is often a function of distinct habitat changes (Geffen et al. 2004), presence of dispersal corridors (Ernest et al. 2003, Geffen et al. 2007), or landscape location (e.g., core vs. peripheral populations; Schwartz et al. 2003). Although the region was intersected by United States interstate highways 10 and 20, neither presented the combination of automobile traffic volume or physical barriers to animal movements observed to reduce gene flow in medium-sized carnivores (Riley et al. 2006).

At the landscape scale, Texas gray fox were continuously distributed, with no discernable gaps in populations. It is likely that the weak genetic structure and pattern of spatial autocorrelation that we observed in the gray fox ORV zone were the result of high rates of dispersal. The western Texas landscape apparently presented no physical barriers to gray fox movements. The genetic data were consistent with recent field observations derived from radiotelemetry and tag returns, which have indicated that gray foxes in the ORV zone may move >20 km from their initial capture site (B. Mesenbrink, USDA, Wildlife Services, National Wildlife Research Center, unpublished data). Historical reports also note occasional long-distance (≤ 80 km) movements in the eastern United States (e.g., Sheldon 1953). Our data suggested that long-distance movements in Texas gray fox (on the order of tens of kilometers) may be more common than previously suspected.

A high rate of dispersal appears a likely explanation for the pattern of population structuring observed in the gray fox ORV zone. Unfortunately, we were unable to perform focused tests for sex-bias in dispersal based on differences in F_{ST} and corrected assignment indices (e.g., v_{AI} ; Goudet et al. 2002). This was because simulated data sets suggest that statistical power for the dispersal tests is not expected to be high if the number of microsatellite loci is < 8 and if the dispersal bias is not pronounced (e.g., $< 80:20$; Goudet et al. 2002). When fixation statistics over the entire data set were compared for biparentally inherited microsatellite markers and maternally inherited mtDNA sequence data, there was little evidence for dispersal bias; fixation indices based on mtDNA did not differ from zero, and only a slight departure was observed from microsatellite markers. We can only conclude that we did not detect firm evidence for sex bias in dispersal in our data set. Further investigation is required to fully establish the existence or prevalence of sex-biased dispersal in gray fox from this region of Texas. Additional hypotheses regarding dispersal that merit investigation include determining if evidence exists for joint

dispersal after pair formation or if movements are variable depending on availability of territories or resources.

MANAGEMENT IMPLICATIONS

Gray fox populations appeared weakly structured and genetically non-independent across a broad (≥ 30 -km) geographic extent, suggesting high rates of movement or dispersal. Thus, a population (assuming a 30-km diam) could conservatively span approximately 707 km², and perhaps farther. Therefore, the historical 16-km to 24-km wide ORV zone barrier (Sidwa et al. 2005) may be too narrow to contain the TF rabies epizootic. Although rabies-positive gray fox have been periodically found outside of the ORV zone, it is uncertain if gray fox incubating rabies virus have actually breached the ORV zone. Plausible alternative hypotheses for the occurrence of rabid individuals outside the ORV zone include persistent, low-level enzootic rabies outside of this zone or interspecific trafficking of the virus. Neither has been detected to date during intensive monitoring following the annual bait distribution campaigns, but these alternatives cannot be conclusively ruled out. Nevertheless, based on our data, we conclude that a spatially extensive ORV zone would be prudent and may in fact be necessary for the effective control of gray fox rabies in Texas. The ORV zone has been expanded in recent years, in part based on preliminary data derived from this study; we recommend that ORV zones for gray fox continue to be managed as spatially extensive. The specific means chosen to implement and maintain a wider ORV zone will ultimately depend upon the resources available and the priorities of the program (e.g., keeping rabies from areas of dense human populations, interspecific variant transmission, etc.). Nonetheless, the width of gray fox ORV zones may be critical to achieve the long-term goal of eliminating the TF variant from the United States.

Managers have begun to recognize that the efficiency and effectiveness of management efforts can be improved through exploiting (Hampton et al. 2004, Robertson and Gemmell 2004) or minimizing population structuring (Epps et al. 2007). At the landscape scale, management units that are defined too conservatively result in wasted effort, whereas failure to control the entire local population may render management ineffective. In this case, the main useful information derived was the detection of weak structure at the landscape scale and the estimation of its spatial extent. This study serves as an example of how genetic tools can assist and focus large-scale wildlife management programs.

ACKNOWLEDGMENTS

We are indebted and offer our sincere thanks to the Texas Department of State Health Services and Texas Wildlife Services for aid with sample collection and logistical assistance, especially B. Sims, J. Sandoval, E. Miles, J. Dorsett, and M. Mapston. In addition, we express our gratitude to the many private and public land stewards who have graciously given access to their lands to conduct this work. This study would not have been possible without the assistance of a great number of individuals from the aforementioned institutions. Funding for this work was provided by the United States

Department of Agriculture, Animal Plant Health Inspection Service, Wildlife Services National Wildlife Research Center and the Jack Berryman Institute. A. Zamorano was supported by a National Science Foundation Undergraduate Mentoring in Environmental Biology program for promoting undergraduate student research. This is contribution 2009-118 of the Caesar Kleberg Wildlife Research Institute.

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Associate Editor: Latch.